		Case 3:08-cv-01748-CRB [	Document 7	Filed 0	5/15/2008	Page 1 of 23
Gordon & Rees, LLP 275 Battery Street, Suite 2000 San Francisco, CA 94111	1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28		N: 146904) P 4200 m CORPORATION IITED STATES THERN DISTRI SAN FRANCI CTRA TICES AND GATION	DISTRI ICT OF SCO DI ) ) ) ) ) ) ) )	CALIFORNI VISION  MDL Docke CASE NO.  PFIZER IN CORPORA SEARLE L COMPLAI	et No. 1699 3:08-cv-1748-CRB IC., PHARMACIA TION, AND G.D. LC'S ANSWER TO
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ANSWER TO COMPLAINT – 3:08-cv-1748-CRB

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NOW COME Defendants Pfizer Inc. (improperly captioned in Plaintiff's Complaint as "Pfizer, Inc.") ("Pfizer"), Pharmacia Corporation ("Pharmacia"), and G.D. Searle LLC (improperly captioned in Plaintiff's Complaint as "G.D. Searle, LLC") ("Searle") (collectively "Defendants"), and file this Answer to Plaintiff's Complaint ("Complaint"), and would respectfully show the Court as follows:

I.

#### PRELIMINARY STATEMENT

The Complaint does not state in sufficient detail when Plaintiff was prescribed or used Bextra® (valdecoxib) ("Bextra®"). Accordingly, this Answer can only be drafted generally. Defendants may seek leave to amend this Answer when discovery reveals the specific time periods in which Plaintiff was prescribed and used Bextra®.

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#### **ANSWER**

Defendants admit that Plaintiff brought this civil action seeking monetary damages, but deny that Plaintiff is entitled to any relief or damages. Defendants admit that, during certain periods of time, Pfizer and Pharmacia marketed and co-promoted Bextra® in the United States to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants admit that, during certain periods of time, Bextra® was manufactured and packaged for Searle, which developed, tested, marketed, co-promoted and distributed Bextra® in the United States to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDA-approved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants deny any wrongful conduct, deny that Bextra® caused Plaintiff injury or damages, and deny the remaining allegations in this paragraph of the Complaint.

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#### **Response to Allegations Regarding Parties**

- Defendants are without knowledge or information sufficient to form a belief as to the 2. truth of the allegations in this paragraph of the Complaint regarding Plaintiff's age and citizenship, and, therefore, deny the same. Defendants deny the remaining allegations in this paragraph of the Complaint.
- 3. Defendants admit that Pfizer is a Delaware corporation with its principal place of business in New York. Defendants admit that Pharmacia acquired Searle in 2000 and that, as the result of a merger in April 2003, Searle and Pharmacia became subsidiaries of Pfizer. Defendants admit that, during certain periods of time, Pfizer marketed and co-promoted Bextra® in the United States, including California and Mississippi, to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants state that Plaintiff's allegations regarding "predecessors in interest" are vague and ambiguous. Defendants are without knowledge or information to form a belief as to the truth of such allegations, and, therefore, deny the same. Defendants deny the remaining allegations in this paragraph of the Complaint.
- Defendants admit that Searle is a Delaware limited liability company with its principal place of business in Illinois. Defendants admit that, as the result of a merger in April 2003, Searle became a subsidiary of Pfizer. Defendants admit that, during certain periods of time, Bextra® was manufactured and packaged for Searle, which developed, tested, marketed, copromoted and distributed Bextra® in the United States to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants deny the remaining allegations in this paragraph of the Complaint.
- Defendants admit that Pharmacia is a Delaware corporation with its principal place of business in New Jersey. Defendants admit that, during certain periods of time, Pharmacia marketed and co-promoted Bextra® in the United States, including California and Mississippi, to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants state that Plaintiff's allegations regarding "predecessors in interest" are vague and ambiguous. Defendants are without

# Response to Allegations Regarding Jurisdiction and Venue

knowledge or information to form a belief as to the truth of such allegations, and, therefore,

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6. Defendants are without knowledge or information to form a belief as to the truth of the allegations in this paragraph of the Complaint regarding the amount in controversy, and, therefore, deny that the same. However, Defendants admit that Plaintiff claims that the amount in controversy exceeds \$75,000, exclusive of interests and costs.

- Defendants are without knowledge or information sufficient to form a belief as to the truth of the allegations in this paragraph of the Complaint regarding Plaintiff's citizenship and the amount in controversy, and, therefore, deny the same. However, Defendants admit that Plaintiff claims that the parties are diverse and that the amount in controversy exceeds \$75,000, exclusive of interests and costs.
- 8. Defendants are without knowledge or information sufficient to form a belief as to the truth of the allegations in this paragraph of the Complaint regarding the judicial district in which the asserted claims allegedly arose, and, therefore, deny the same. Defendants deny any wrongful conduct, deny committing a tort in the State of California, and deny the remaining allegations in this paragraph of the Complaint.
- 9. Defendants admit that, during certain periods of time, Pfizer and Pharmacia marketed and co-promoted Bextra® in the United States, including California, to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants admit that, during certain periods of time, Bextra® was manufactured and packaged for Searle, which developed, tested, marketed, co-promoted and distributed Bextra® in the United States to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants admit that they provided FDA-approved prescribing information regarding Bextra®. Defendants admit that they do business in the State of California. Defendants state that Plaintiff's allegations regarding "predecessors in interest" are vague and ambiguous. Defendants are without knowledge or information to form a belief as to the truth of such

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allegations, and, therefore, deny the same. Defendants deny any wrongful conduct and deny the remaining allegations in this paragraph of the Complaint.

#### Response to Allegations Regarding Interdistrict Assignment

10. Defendants state that this paragraph of the Complaint contains legal contentions to which no response is required. To the extent that a response is deemed required, Defendants admit that this case should be transferred to In re: Bextra and Celebrex Marketing, Sales Prac. and Prods. Liab. Litig., MDL-1699, assigned to the Honorable Charles R. Breyer by the Judicial Panel on Multidistrict Litigation on September 6, 2005.

#### **Response to Factual Allegations**

- Defendants admit that, during certain periods of time, Pfizer and Pharmacia marketed 11. and co-promoted Bextra® in the United States, including Arkansas, to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants admit that, during certain periods of time, Bextra® was manufactured and packaged for Searle, which developed, tested, marketed, co-promoted and distributed Bextra® in the United States to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants admit that they provided FDA-approved prescribing information regarding Bextra®. Defendants deny the remaining allegations in this paragraph of the Complaint.
- 12. Defendants are without knowledge or information sufficient to form a belief as to the truth of the allegations in this paragraph of the Complaint regarding whether Plaintiff used Bextra®, and, therefore, deny the same. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants deny any wrongful conduct, deny that Bextra® caused Plaintiff injury or damages, and deny the remaining allegations in this paragraph of the Complaint.
- 13. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDA-approved prescribing information, which was at all times adequate and comported with applicable standards of care and law.

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- Defendants admit that, during certain periods of time, Pfizer and Pharmacia marketed and co-promoted Bextra® in the United States to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants admit that, during certain periods of time, Bextra® was manufactured and packaged for Searle, which developed, tested, marketed, co-promoted and distributed Bextra® in the United States to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants admit that they provided FDAapproved prescribing information regarding Bextra®. Defendants deny the remaining allegations in this paragraph of the Complaint.
- Defendants admit that Bextra® is in a class of drugs that is, at times, referred to as non-15. steroidal anti-inflammatory drugs ("NSAIDS"). Defendant states that, as stated in the FDAapproved labeling for Bextra®, "[t]he mechanism of action is believed to be due to inhibition of prostaglandin synthesis primarily through inhibition of cyclooxygenase-2 (COX-2). therapeutic plasma concentrations in humans valdecoxib does not inhibit cyclooxygenase-1 (COX-1)." Defendants admit that Bextra® was approved by the FDA on November 16, 2001. Defendants admit, as indicated in the package insert approved by the FDA, that Bextra® is indicated for use in the relief of the signs and symptoms of osteoarthritis and adult rheumatoid arthritis, as well as for the treatment of primary dysmenorrhea. Defendants deny the remaining allegations in this paragraph of the Complaint.
- 16. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDA-approved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants deny any wrongful conduct and deny the remaining the allegations in this paragraph of the Complaint.
- 17. Defendants are without knowledge or information sufficient to form a belief as to the truth of the allegations in this paragraph of the Complaint regarding whether Plaintiff used

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Bextra®, and, therefore, deny the same. Defendants state that Bextra® was and is safe and
effective when used in accordance with its FDA-approved prescribing information. Defendants
state that the potential effects of Bextra® were and are adequately described in its FDA-
approved prescribing information, which was at all times adequate and comported with
applicable standards of care and law. Defendants deny any wrongful conduct and deny the
remaining the allegations in this paragraph of the Complaint.

- Defendants state that Bextra® was and is safe and effective when used in accordance 18. with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDA-approved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants admit that they provided FDA-approved prescribing information regarding Bextra®. Defendants deny the remaining the allegations in this paragraph of the Complaint.
- 19 Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDA-approved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants deny any wrongful conduct and deny the remaining the allegations in this paragraph of the Complaint.
- 20. Defendants are without knowledge or information sufficient to form a belief as to the truth of the allegations in this paragraph of the Complaint regarding whether Plaintiff used Bextra®, and, therefore, deny the same. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDAapproved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants deny any wrongful conduct, deny that Bextra® caused Plaintiff injury or damages, and deny the remaining the allegations in this paragraph of the Complaint.

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#### **Response to First Cause of Action: Products Liability**

- 21. Defendants incorporate by reference their responses to each paragraph of Plaintiff's Complaint as if fully set forth herein.
- 22. Defendants admit that, during certain periods of time, Pfizer and Pharmacia marketed and co-promoted Bextra® in the United States, including California, to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants admit that, during certain periods of time, Bextra® was manufactured and packaged for Searle, which developed, tested, marketed, co-promoted and distributed Bextra® in the United States to be prescribed by healthcare providers who are by law authorized to prescribe drugs in accordance with their approval by the FDA. Defendants admit that they provided FDA-approved prescribing information regarding Bextra®. Defendants deny the remaining the allegations in this paragraph of the Complaint.
- 23. Defendants admit that Bextra® was expected to reach consumers without substantial change from the time of sale. Defendants are without knowledge or information sufficient to form a belief as to the truth of the allegations in this paragraph of the Complaint regarding whether Plaintiff used Bextra®, and, therefore, deny the same. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDA-approved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants deny any wrongful conduct, deny that Bextra® is unreasonably dangerous, and deny the remaining allegations this paragraph of the Complaint.
- 24. Defendants are without knowledge or information sufficient to form a belief as to the truth of the allegations in this paragraph of the Complaint regarding whether Plaintiff used Bextra®, and, therefore, deny the same. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDAapproved prescribing information, which was at all times adequate and comported with

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Gordon & Rees, LLP 275 Battery Street, Suite 2000 San Francisco, CA 94111 applicable standards of care and law. Defendants deny any wrongful conduct, deny that Bextra® is defective or unreasonably dangerous, deny that Bextra® caused Plaintiff injury or damages, and deny the remaining allegations this paragraph of the Complaint.

- 25. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDA-approved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants deny any wrongful conduct, deny that Bextra® caused Plaintiff injury or damages, and deny the remaining allegations this paragraph of the Complaint.
- Defendants state that Bextra® was and is safe and effective when used in accordance 26. with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDA-approved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants deny any wrongful conduct and deny the remaining allegations this paragraph of the Complaint.
- 27. Defendants are without knowledge or information sufficient to form a belief as to the truth of the allegations in this paragraph of the Complaint regarding whether Plaintiff used Bextra®, and, therefore, deny the same. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDAapproved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants deny any wrongful conduct and deny the remaining allegations this paragraph of the Complaint.
- 28. Defendants are without knowledge or information sufficient to form a belief as to the truth of the allegations in this paragraph of the Complaint regarding whether Plaintiff used Bextra®, and, therefore, deny the same. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDA-

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- approved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants deny any wrongful conduct, deny that Bextra® caused Plaintiff injury or damages, and deny the remaining allegations this paragraph of the Complaint.
- 29. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDA-approved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants deny any wrongful conduct and deny the remaining allegations this paragraph of the Complaint.
- 30. Defendants state that Bextra® was and is safe and effective when used in accordance with its FDA-approved prescribing information. Defendants state that the potential effects of Bextra® were and are adequately described in its FDA-approved prescribing information, which was at all times adequate and comported with applicable standards of care and law. Defendants deny any wrongful conduct, deny that Bextra® caused Plaintiff injury or damages, and deny the remaining allegations this paragraph of the Complaint.
- 31. Defendants deny any wrongful conduct, deny that Bextra® caused Plaintiff injury or damages, and deny the remaining allegations this paragraph of the Complaint.

# Response to Allegations Regarding Damages

- 32. Defendants incorporate by reference their responses to each paragraph of Plaintiff's Complaint as if fully set forth herein.
- Defendants deny any wrongful conduct, deny that Bextra® caused Plaintiff injury or 33. damages, and deny the remaining allegations this paragraph of the Complaint, including all subparts.
- 34. Answering the first unnumbered paragraph following Paragraph 33 of the Complaint, Defendants deny any wrongful conduct, deny that Bextra® caused Plaintiff injury or damages, and deny the remaining allegations this paragraph of the Complaint.
- 35. Answering the second unnumbered paragraph following Paragraph 33 of the Complaint,

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Defendants deny any wrongful conduct, deny that Bextra® caused Plaintiff injury or damages, and deny the remaining allegations this paragraph of the Complaint, including all subparts.

III.

#### **GENERAL DENIAL**

Defendants deny all allegations and/or legal conclusions set forth in Plaintiff's Complaint that have not been previously admitted, denied, or explained.

IV.

#### **AFFIRMATIVE DEFENSES**

Defendants reserve the right to rely upon any of the following or additional defenses to claims asserted by Plaintiff to the extent that such defenses are supported by information developed through discovery or evidence at trial. Defendants affirmatively show that:

#### **First Defense**

1. The Complaint fails to state a claim upon which relief can be granted.

#### **Second Defense**

2. Bextra® is a prescription medical product. The federal government has preempted the field of law applicable to the labeling and warning of prescription medical products. Defendants' labeling and warning of Bextra® was at all times in compliance with applicable federal law. Plaintiff's causes of action against Defendants, therefore, fail to state a claim upon which relief can be granted; such claims, if allowed, would conflict with applicable federal law and violate the Supremacy Clause of the United States Constitution.

#### **Third Defense**

3. At all relevant times, Defendants provided proper warnings, information and instructions for the drug in accordance with generally recognized and prevailing standards in existence at the time.

#### **Fourth Defense**

4. At all relevant times, Defendants' warnings and instructions with respect to the use of Bextra® conformed to the generally recognized, reasonably available, and reliable state of knowledge at the time the drug was manufactured, marketed and distributed.

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Fifth	Defense
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5. Plaintiff's action is time-barred as it is filed outside of the time permitted by the applicable Statute of Limitations, and same is pled in full bar of any liability as to Defendants.

#### **Sixth Defense**

6. Plaintiff's action is barred by the statute of repose.

#### **Seventh Defense**

7. If Plaintiff sustained any injuries or incurred any losses or damages as alleged in the Complaint, the same were caused by the negligence or fault of the Plaintiff and Plaintiff's damages, if any, are barred or reduced by the doctrines of comparative fault and contributory negligence and by the failure to mitigate damages.

#### **Eighth Defense**

8. The proximate cause of the loss complained of by Plaintiff is not due to any acts or omissions on the part of Defendants. Rather, said loss is due to the acts or omissions on the part of third parties unrelated to Defendants and for whose acts or omissions Defendants are not liable in any way.

#### Ninth Defense

9. The acts and/or omissions of unrelated third parties as alleged constituted independent, intervening causes for which Defendants cannot be liable.

#### **Tenth Defense**

10. Any injuries or expenses incurred by Plaintiff were not caused by Bextra®, but were proximately caused, in whole or in part, by an idiosyncratic reaction, operation of nature, or act of God.

#### **Eleventh Defense**

11. Defendants affirmatively deny that they violated any duty owed to Plaintiff.

#### **Twelfth Defense**

12. A manufacturer has no duty to warn patients or the general public of any risk, contraindication, or adverse effect associated with the use of a prescription medical product. Rather, the law requires that all such warnings and appropriate information be given to the

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prescribing physician and the medical profession, which act as a "learned intermediary" in determining the use of the product. Bextra® is a prescription medical product, available only on the order of a licensed physician. Bextra® provided an adequate warning to Plaintiff's treating and prescribing physicians.

#### **Thirteenth Defense**

13. The product at issue was not in a defective condition or unreasonably dangerous at the time it left the control of the manufacturer or seller.

#### **Fourteenth Defense**

14. Bextra® was at all times material to the Complaint reasonably safe and reasonably fit for its intended use and the warnings and instructions accompanying Bextra® at the time of the occurrence of the injuries alleged by Plaintiff were legally adequate for its approved usages.

#### **Fifteenth Defense**

15. Plaintiff's causes of action are barred in whole or in part by the lack of a defect as the Bextra® allegedly ingested by Plaintiff was prepared in accordance with the applicable standard of care.

#### Sixteenth Defense

16. If Plaintiff sustained any injuries or incurred any losses or damages as alleged in the Complaint, the same were caused by the unforeseeable alteration, change, improper handling, abnormal use, or other unforeseeable misuse of Bextra® by persons other than Defendants or persons acting on its behalf after the product left the control of Defendants.

#### **Seventeenth Defense**

Plaintiff's alleged damages were not caused by any failure to warn on the part of 17. Defendants.

#### **Eighteenth Defense**

18. Plaintiff's alleged injuries/damages, if any, were the result of preexisting or subsequent conditions unrelated to Bextra®.

#### **Nineteenth Defense**

19. Plaintiff knew or should have known of any risk associated with Bextra®; therefore, the San Francisco, CA 94111

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doctrine of assumption of the risk bars or diminishes any recovery.

#### **Twentieth Defense**

20. Plaintiff is barred from recovering against Defendants because Plaintiff's claims are preempted in accordance with the Supremacy Clause of the United States Constitution and by the Federal Food, Drug and Cosmetics Act, 21 U.S.C. § 301 et. seg.

#### **Twenty-first Defense**

21. Plaintiff's claims are barred in whole or in part under the applicable state law because the subject pharmaceutical product at issue was subject to and received pre-market approval by the Food and Drug Administration under 52 Stat. 1040, 21 U.S.C. § 301.

#### **Twenty-second Defense**

The manufacture, distribution and sale of the pharmaceutical product referred to in 22. Plaintiff's Complaint were at all times in compliance with all federal regulations and statutes, and Plaintiff's causes of action are preempted.

#### **Twenty-third Defense**

Plaintiff's claims are barred in whole or in part by the deference given to the primary 23. jurisdiction of the Food and Drug Administration over the subject pharmaceutical product at issue under applicable federal laws, regulations, and rules.

#### **Twenty-fourth Defense**

24. Plaintiff's claims are barred in whole or in part because there is no private right of action concerning matters regulated by the Food and Drug Administration under applicable federal laws, regulations, and rules.

#### **Twenty-fifth Defense**

25. Plaintiff's claims are barred in whole or in part because Defendants provided adequate "direction or warnings" as to the use of the subject pharmaceutical product within the meaning of Comment j to Section 402A of the Restatement (Second) of Torts.

#### **Twenty-sixth Defense**

26. Plaintiff's claims are barred or limited to a product liability failure to warn claim because Bextra® is a prescription pharmaceutical drug and falls within the ambit of

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Gordon & Rees, LLP 275 Battery Street, Suite 2000 Restatement (Second) of Torts § 402A, Comment k.

#### **Twenty-seventh Defense**

27. Plaintiff's claims are barred in whole or in part because the subject pharmaceutical product at issue "provides net benefits for a class of patients" within the meaning of Comment f to § 6 of the Restatement (Third) of Torts: Products Liability.

#### **Twenty-eighth Defense**

28. Plaintiff's claims are barred under § 4, et seq., of the Restatement (Third) of Torts: Products Liability.

#### **Twenty-ninth Defense**

29. To the extent that Plaintiff is seeking punitive damages, Plaintiff has failed to plead facts sufficient under the law to justify an award of punitive damages.

#### **Thirtieth Defense**

30. The imposition of punitive damages in this case would violate Defendants' rights to procedural due process under the Fourteenth Amendment of the United States Constitution and the Constitution of the States of California and Arkansas, and would additionally violate Defendants' right to substantive due process under the Fourteenth Amendment of the United States Constitution.

#### **Thirty-first Defense**

31. Plaintiff's claims for punitive damages are barred, in whole or in part, by the Fifth and Fourteenth Amendments to the United States Constitution.

#### **Thirty-second Defense**

32. The imposition of punitive damages in this case would violate the First Amendment to the United States Constitution.

#### **Thirty-third Defense**

33. Plaintiff's punitive damage claims are preempted by federal law.

#### **Thirty-fourth Defense**

34. In the event that reliance was placed upon Defendants' nonconformance to an express representation, this action is barred as there was no reliance upon representations, if any, of

# Defendants.

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#### 35. Plaintiff failed to provide Defendants with timely notice of any alleged nonconformance

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to any express representation.

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# **Thirty-sixth Defense**

**Thirty-fifth Defense** 

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# **Thirty-seventh Defense**

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labeling with respect to the subject pharmaceutical products were not false or misleading and,

Plaintiff's claims are barred, in whole or in part, because the advertisements, if any, and

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therefore, constitute protected commercial speech under the applicable provisions of the United

States Constitution.

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#### **Thirty-eighth Defense**

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38. To the extent that Plaintiff seeks punitive damages for the conduct which allegedly caused injuries asserted in the Complaint, punitive damages are barred or reduced by applicable law or statute or, in the alternative, are unconstitutional insofar as they violate the due process protections afforded by the United States Constitution, the excessive fines clause of the Eighth Amendment of the United States Constitution, the Commerce Clause of the United States Constitution, and the Full Faith and Credit Clause of the United States Constitution and the Constitution of the States of California and Arkansas. Any law, statute, or other authority purporting to permit the recovery of punitive damages in this case is unconstitutional, facially and as applied, to the extent that, without limitation, it: (1) lacks constitutionally sufficient standards to guide and restrain the jury's discretion in determining whether to award punitive damages and/or the amount, if any; (2) is void for vagueness in that it failed to provide adequate advance notice as to what conduct will result in punitive damages; (3) permits recovery of punitive damages based on out-of-state conduct, conduct that complied with applicable law, or

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conduct that was not directed, or did not proximately cause harm, to Plaintiff; (4) permits

recovery of punitive damages in an amount that is not both reasonable and proportionate to the

538 U.S. 408 (2003).

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permits jury consideration of net worth or other financial information relating to Defendants; (6) lacks constitutionally sufficient standards to be applied by the trial court in post-verdict review of any punitive damages awards; (7) lacks constitutionally sufficient standards for appellate review of punitive damages awards; and (8) otherwise fails to satisfy Supreme Court precedent, including, without limitation, Pacific Mutual Life Ins. Co. v. Haslip, 499 U.S. 1

(1991), TXO Production Corp. v. Alliance Resources, Inc., 509 U.S. 443 (1993); BMW of North

amount of harm, if any, to Plaintiff and to the amount of compensatory damages, if any; (5)

America, Inc. v. Gore, 519 U.S. 559 (1996); and State Farm Mut. Auto Ins. Co. v. Campbell,

# **Thirty-ninth Defense**

39. The methods, standards, and techniques utilized with respect to the manufacture, design, and marketing of Bextra®, if any, used in this case, included adequate warnings and instructions with respect to the product's use in the package insert and other literature, and conformed to the generally recognized, reasonably available, and reliable state of the knowledge at the time the product was marketed.

# **Fortieth Defense**

40. The claims asserted in the Complaint are barred because Bextra® was designed, tested, manufactured and labeled in accordance with the state-of-the-art industry standards existing at the time of the sale.

#### **Forty-first Defense**

41. If Plaintiff has sustained injuries or losses as alleged in the Complaint, upon information and belief, such injuries and losses were caused by the actions of persons not having real or apparent authority to take said actions on behalf of Defendants and over whom Defendants had no control and for whom Defendants may not be held accountable.

#### **Forty-second Defense**

42. The claims asserted in the Complaint are barred, in whole or in part, because Bextra® was not unreasonably dangerous or defective, was suitable for the purpose for which it was intended, and was distributed with adequate and sufficient warnings.

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Forty	v-third	Defense
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43. Plaintiff's claims are barred, in whole or in part, by the equitable doctrines of laches, waiver, and/or estoppel.

#### **Forty-fourth Defense**

44. Plaintiff's claims are barred because Plaintiff's injuries, if any, were the result of the pre-existing and/or unrelated medical, genetic and/or environmental conditions, diseases or illnesses, subsequent medical conditions or natural courses of conditions of Plaintiff, and were independent of or far removed from Defendants' conduct.

#### **Forty-fifth Defense**

45. The claims asserted in the Complaint are barred, in whole or in part, because Bextra® did not proximately cause injuries or damages to Plaintiff.

#### **Forty-sixth Defense**

46. The claims asserted in the Complaint are barred, in whole or in part, because Plaintiff did not incur any ascertainable loss as a result of Defendants' conduct.

#### **Forty-seventh Defense**

47. The claims asserted in the Complaint are barred, in whole or in part, because the manufacturing, labeling, packaging, and any advertising of the product complied with the applicable codes, standards and regulations established, adopted, promulgated or approved by any applicable regulatory body, including but not limited to the United States, any state, and any agency thereof.

#### **Forty-eighth Defense**

The claims must be dismissed because Plaintiff would have taken Bextra® even if the 48. product labeling contained the information that Plaintiff contends should have been provided.

#### **Forty-ninth Defense**

49. The claims asserted in the Complaint are barred because the utility of Bextra® outweighed its risks.

#### Fiftieth Defense

Plaintiff's damages, if any, are barred or limited by the payments received from 50.

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#### **Fifty-first Defense**

51. Defendants' liability, if any, can only be determined after the percentages of responsibility of all persons who caused or contributed toward Plaintiff's alleged damages, if any, are determined. Defendants seek an adjudication of the percentage of fault of the claimants and each and every other person whose fault could have contributed to the alleged injuries and damages, if any, of Plaintiff.

#### **Fifty-second Defense**

52. Plaintiff's claims are barred, in whole or in part, by the doctrine of abstention in that the common law gives deference to discretionary actions by the United States Food and Drug Administration under the Federal Food, Drug, and Cosmetic Act.

#### **Fifty-third Defense**

53. The claims asserted in the Complaint are barred, in whole or in part, because Bextra® is comprehensively regulated by the FDA pursuant to the Federal Food, Drug & Cosmetic Act ("FDCA"), 21 U.S.C. §§ 301 et seq., and regulations promulgated there under, and Plaintiff's claims conflict with the FDCA, with the regulations promulgated by FDA to implement the FDCA, with the purposes and objectives of the FDCA and FDA's implementing regulations, and with the specific determinations by FDA specifying the language that should be used in the labeling accompanying Bextra®. Accordingly, Plaintiff's claims are preempted by the Supremacy Clause of the United States Constitution, Article VI, clause 2, and the laws of the United States.

# **Fifty-fourth Defense**

54. Plaintiff's misrepresentation allegations are not stated with the degree of particularity required by Federal Rule of Civil Procedure 9(b) and should be dismissed.

#### **Fifty-fifth Defense**

55. Defendants state on information and belief that the Complaint and each purported cause of action contained therein is barred by the statutes of limitations contained in California Code of Civil Procedure §§ 335.1 and 338 and former § 340(3), such other statutes of limitation as may apply.

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#### **Fifty-sixth Defense**

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56. Defendants state on information and belief that any injuries, losses, or damages suffered by Plaintiff were proximately caused, in whole or in part, by the negligence or other actionable

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conduct of persons or entities other than Defendants. Therefore, Plaintiff's recovery against Defendants, if any, should be reduced pursuant to California Civil Code § 1431.2. **Fifty-seventh Defense** 

57. To the extent that Plaintiff seeks punitive damages for an alleged act or omission of Defendants, no act or omission was oppressive, fraudulent, or malicious under California Civil Code § 3294, and, therefore, any award of punitive damages is barred. Any claim for punitive damages is also barred under California Civil Code § 3294(b).

Fifty-eighth Defense

Plaintiff's fraud based claims, if any, are not stated with particularity as required by 58. Rule 9 of the Arkansas Rules of Civil Procedure.

**Fifty-ninth Defense** 

Plaintiff's damages, if any, must be reduced by the percentage of fault attributable to 59. Plaintiff and to nonparties as provided by Ark. Code Ann. § 16-55-202.

**Sixtieth Defense** 

60. Plaintiff's claims are barred and/or limited by the provisions of the Arkansas Products Liability Act, Ark. Code Ann. § 16-116-101, et seq.

**Sixty-first Defense** 

61. Plaintiff's claims for punitive damages are barred, in whole or in part, by the Arkansas Civil Justice Reform Act of 2003, Ark. Code Ann. § 16-55-201, et seq.

**Sixty-second Defense** 

62. Defendants reserve the right to supplement their assertion of defenses as they continue with their factual investigation of Plaintiff's claims.

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	q	ase 3:08-cv-01748-CRB Document 7 Filed 05/15/2008 Page 21 of 23				
	1	V.				
	2	<u>PRAYER</u>				
	3	WHEREFORE, Defendants pray for judgment as follows:				
	4	1. That Plaintiff takes nothing from Defendants by reason of the Complaint;				
	5	2. That the Complaint be dismissed;				
	6	3. That Defendants be awarded their costs for this lawsuit;				
	7	4. That the trier of fact determine what percentage of the combined fault or other liability				
	8	of all persons whose fault or other liability proximately caused Plaintiff's alleged				
	9	injuries, losses or damages is attributable to each person;				
	10	5. That any judgment for damages against Defendants in favor of Plaintiff be no greater				
	11	than an amount which equals their proportionate share, if any, of the total fault or other				
LLP nite 2000 94111	12	liability which proximately caused Plaintiff's injuries and damages; and				
Gordon & Rees, LLP 5 Battery Street, Suite 2000 San Francisco, CA 94111	13	6. That Defendants have such other and further relief as the Court deems appropriate.				
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ANSWER TO COMPLAINT – 3:08-cv-1748-CRB

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275 S	17			CORPORATI LLC	ON, AND G.D. SEARLE
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ANSWER TO COMPLAINT – 3:08-cv-1748-CRB

#### 1 **JURY DEMAND** 2 Defendants Pfizer Inc., Pharmacia Corporation, and G.D. Searle LLC, hereby demand a 3 trial by jury of all the facts and issues in this case pursuant to 38(b) of the Federal Rules of Civil 4 Procedure. 5 May 15, 2008 GORDON & REES LLP 6 7 By:\_ 8 Stuart M. Gordon sgordon@gordonrees.com Embarcadero Center West 275 Battery Street, 20<sup>th</sup> Floor San Francisco, CA 94111 9 10 Telephone: (415) 986-5900 11 Fax: (415) 986-8054 275 Battery Street, Suite 2000 12 San Francisco, CA 94111 Gordon & Rees, LLP May 15, 2008 TUCKER ELLIS & WEST LLP 13 14 By:\_\_\_\_\_ 15 Michael C. Zellers 16 michael.zellers@tuckerellis.com 515 South Flower Street, Suite 4200 17 Los Angeles, CA 90071-2223 Telephone: (213) 430-3400 18 Fax: (213) 430-3409 19 Attorneys for Defendants PFIZER INC., PHARMACIA 20 CORPORATION, AND G.D. SEARLE LLC 21 22 23 24 25 26 27 28